



(1) Publication number:

0 540 009 A1

12

# **EUROPEAN PATENT APPLICATION**

(1) Application number: 92118554.2

(f) Int. Cl.5: **C07D** 301/06, C07D 303/02

2 Date of filing: 29.10.92

Priority: 30.10.91 JP 284651/91 11.03.92 JP 52438/92 11.03.92 JP 52441/92

② Date of publication of application: 05.05.93 BulletIn 93/18

Designated Contracting States:
 BE CH DE FR GB IT LI NL

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Process for producing epoxides.

A process for producing an epoxide represented by the formula (2) which comprises reacting an olefin represented by the formula (1) with oxygen in the presence of an aldehyde and in the presence or absence of a proton source by using no catalyst or using an iron – containing or copper – containing catalyst.

$$C = C$$

$$R^{2}$$

$$R^{2}$$

 $R^1$  C - C

Formula (1)

Formula (2)

The present invention relates to a process for producing an epoxide represented by the formula (2) shown later.

The epoxide mentioned above is important as an intermediate in producing a wide variety of products, including perfumes, pharmaceuticals, agricultural chemicals, liquid crystals and resins.

It is well known to convert an olefin to the corresponding epoxide by using peracids such as peracetic acid, m-chloroperbenzoic acid, and the like (Some Modern Methods of Organic Synthesis, 3rd ed., P. 370-373). However, since peracids are highly sensitive to shock and are explosive, this method cannot be said to be an advantageous process from the industrial point of view.

To overcome such difficulty, processes have already been developed which comprise oxidizing an olefin with oxygen in the presence of an aldehyde by using a catalyst containing a soluble praseodymium compound (Japanese Patent Application KOKAI No. 59 – 231077) or a soluble nickel catalyst (Chem. Lett., 1991, 1). However, since these processes use a homogeneous catalyst, they require complicated operations in the separation of the epoxide, the intended product, from the catalyst and in the recovery of the catalyst, and further, when the catalyst is not recovered, the processes will produce a waste water with a high content of undesirable substances. Thus, they are not fully satisfactory as industrial processes.

The object of the present invention is to provide an advantageous process for producing epoxides by oxidizing olefins with oxygen in the presence of an aldehyde which uses no catalyst or uses a catalyst that in easily available and recoverable and that will bring little of undesirable substances into the waste water.

This object has been attained by the present invention.

Thus, according to the present invention, there is provided a process for producing an epoxide represented by the formula (2)

$$\begin{array}{c|c}
R^1 & O & R^3 \\
C - C & \\
R^2 & R^4
\end{array}$$

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wherein  $R^1$ ,  $R^2$ ,  $R^3$  and  $R^4$  may be the same or different and each denotes a hydrogen atom,  $(C_1 - C_{20})$ alkyl group; alkyl group substituted with a halogen, hydroxy, alkoxy, phenoxy, acyloxy, acyl, alkoxycarbonyl or phenoxycarbonyl; phenyl group substituted with a halogen, alkyl, alkoxy, phenoxy, acyloxy, acyl, alkoxycarbonyl or phenoxycarbonyl; phenylalkyl group, phenylalkyl group substituted with a halogen, alkyl, alkoxy, phenoxy, acyloxy, acyl, alkoxycarbonyl or phenoxycarbonyl; acyl group, alkoxycarbonyl group or phenoxycarbonyl group; provided that  $R^1$  and  $R^2$  or  $R^1$  and  $R^3$  may combine with each other to form a ring, or  $R^1$ ,  $R^2$  and  $R^3$  may combine altogether to form a condensed ring, which comprises reacting an olefin represented by the formula (1)

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wherein R<sup>1</sup>, R<sup>2</sup>, R<sup>3</sup> and R<sup>4</sup> are the same as defined above, with oxygen in the presence of an aldehyde, preferably in the presence or absence of a proton source and in the presence or absence of an iron – containing catalyst or a copper – containing catalyst.

Generally, the terms alkyl and acyl, alone or as part of another group, comprise residues having up to 20, preferably up to 10, more preferably 1 to 6 carbon atoms.

Examples of the olefin represented by the formula (1) used in the present invention include ethylene, propylene, butene, pentene, hexene, heptene, octene, nonene, decene, undecene, dodecene, pentadecene, eicocene, methylbutene, methylpentene, methylhexene, methylhexene, methylhexene, methylhexene, dimethylhexene, dimethylhexene, dimethylhexene, dimethylhexene, trimethylnonene, ethylpentene, ethylhexene, ethylhexene, n-propylnonene, tetramethylnonene, tetramethyldecene, ethyl-

n-propyldecene, cyclopentene, cyclohexene, cyclohexene, cyclooctene, cyclododecene, methyl cyclopentene, methylcyclohexene, ethylcyclohexene, ethylcyclooctene, dimethylcyclohexene, norbornene, pinene, allyl chloride, allyl bromide, crotyl chloride, crotyl bromide, 1,4-dichlorobutene, pentenol, cyclohexenol, terpineol, methyl pentenyl ether, cyclohexenyl ethyl ether, cyclohexenyl phenyl ether, pentenyl acetate, cyclohexenyl acetate, pentenone, hexenone, heptenone, methyl hexenoate, ethyl oleate, phenyl hexenoate, phenyl oleate, styrene, methylstyrene, ethylstyrene, stilbene, p-chlorostyrene, m-chlorostyrene, p-methylstyrene, p-methoxystyrene, p-ethoxystyrene, p-ethoxystyrene, p-methoxystyrene, p-methoxycarbonylstyrene, p-phenoxycarbonylstyrene, p-phenoxycar-phenylbutene, p-methylphenylbutene, p-ethylphenylbutene, p-methoxyphenylbutene, p-ethox-yphenylbutene, p-methoxycarbonylphenylbutene, p-methox-ycarbonylphenylbutene, p-phenoxycarbonylphenylbutene, p-methox-ycarbonylphenylbutene, p-phenoxycarbonylphenylbutene, cyclohexenone, methyl cinnamate, methyl p-methoxycinnamate, phenyl cinnamate, cholesterol, and cholesteryl acetate. The positions of the substituents in the olefins are optional, and the olefins include, if any, the geometric isomers.

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The epoxide which is the intended compound of the present invention can be obtained as the epoxide represented by the formula (2) by using the above-mentioned olefin. Examples of the epoxide include ethylene oxide, propylene oxide, butene oxide, pentene oxide, hexene oxide, heptene oxide, octene oxide, nonene oxide, decene oxide, undecene oxide, dodecene oxide, pentadecene oxide, eicocene oxide, methylbutene oxide, methylpentene oxide, methylhexene oxide, methylhexene oxide, methylbetene oxide, methy methyltetradecene oxide, dimethylbutene oxide, dimethylpentene oxide, dimethylhexene oxide, dimethylheptene oxide, dimethyldecene oxide, trimethylnonene oxide, ethylpentene oxide, ethylpentene oxide, ethylheptene oxide, n - propylnonene oxide, tetramethylnonene oxide, tetramethyldecene oxide, ethyl - n propyldecene oxide, cyclopentene oxide, cyclohexene oxide, cyclohexene oxide, cyclohexene oxide, cyclohexene oxide, cyclododecene oxide, methylcyclopentene oxide, methylcyclohexene oxide, ethylcyclohexene oxide, ethylc cyclooctene oxide, dimethylcyclohexene oxide, norbornene oxide, pinene oxide, epichlorohydrin, epi-1 - chloro - 2,3 - epoxybutane, 1 - bromo - 2,3 - epoxybutane, 1,4 - dichloro - 2,3 - epox ybutane, epoxypentanol, epoxycyclohexanol, terpineol oxide, epoxypentyl methyl ether, epoxycyclohexyl ethyl ether, epoxycyclohexyl phenyl ether, epoxypentyl acetate, epoxycyclohexyl acetate, epoxypentanone, epoxyhexanone, epoxyheptanone, methyl epoxyhexanoate, ethyl epoxyoctadecanoate, phenyl epoxyhexanoate, phenyl epoxyoctadecanoate, styrene oxide, methylstyrene oxide, ethylstyrene oxide, stilbene oxide, p-chlorostyrene oxide, m-chlorostyrene oxide, p-methylstyrene oxide, p-ethylstyrene oxide, pmethoxystyrene oxide, p - ethoxystyrene oxide, 1 - (3',4' - dimethoxyphenyl) - 1,2 - epoxypropane, p - ac etoxystyrene oxide, p - acetylstyrene oxide, p - methoxycarbonylstyrene oxide, p - phenoxycarbonylstyrene oxide, phenylbutene oxide, phenylpentene oxide, phenylhexene oxide, phenyloctene oxide, pchlorophenylbutene oxide, m - chlorophenylbutene oxide, p - methylphenylbutene oxide, p - ethylphenyl butene oxide, p - methoxyphenylbutene oxide, p - ethoxyphenylbutene oxide, m - phenoxyphenylbutene oxide, p - acetoxyphenylbutene oxide, p - acetylphenylbutene oxide, p - methoxycarbonylphenylbutene ox ide, p-phenoxycarbonylphenylbutene oxide, epoxycyclohexanone, methyl 3-phenylglycidate, methyl 3-(4-methoxyphenyl)glycidate, phenyl 3-phenylglycidate, 5,6-epoxy-3-cholestanol and 5,6-epoxy-3cholestanyl acetate.

Examples of the iron – containing catalyst include Fe, Fe(CO)<sub>5</sub>, Fe(CO)<sub>3</sub>, Fe(CO)<sub>12</sub>, FeO, FeCl<sub>2</sub>  $\circ$  nH<sub>2</sub>O, FeBr<sub>2</sub>  $\circ$  nH<sub>2</sub>O, FeSO<sub>4</sub>  $\circ$  nH<sub>2</sub>O, FeCl<sub>3</sub>  $\circ$  nH<sub>2</sub>O, Fe<sub>2</sub>(SO<sub>4</sub>)<sub>3</sub>  $\circ$  nH<sub>2</sub>O, Fe(NO<sub>3</sub>)<sub>3</sub>  $\circ$  nH<sub>2</sub>O, Fe(OAc)<sub>3</sub>, nere preferably Fe, FeCl<sub>2</sub>  $\circ$  nH<sub>2</sub>O, FeSO<sub>4</sub>  $\circ$  nH<sub>2</sub>O, FeCl<sub>3</sub>  $\circ$  nH<sub>2</sub>O, Fe(OAc)<sub>3</sub> and Fe<sub>2</sub>O<sub>3</sub>, more preferably Fe and Fe<sub>2</sub>O<sub>3</sub>. Examples of the copper – containing catalyst include Cu, CuCl, CuCl<sub>2</sub>  $\circ$  nH<sub>2</sub>O, CuBr, CuBr<sub>2</sub>, Cul, CuF<sub>2</sub>, CuSO<sub>4</sub>  $\circ$  nH<sub>2</sub>O, Cu(NO<sub>3</sub>)<sub>3</sub>  $\circ$  nH<sub>2</sub>O, Cu(ClO<sub>4</sub>)<sub>2</sub>  $\circ$  nH<sub>2</sub>O, Cu(OH)<sub>2</sub>, Cu(OCH<sub>3</sub>)<sub>2</sub>, Cu<sub>3</sub>(PO<sub>4</sub>)<sub>2</sub>  $\circ$  nH<sub>2</sub>O, Cu<sub>2</sub>O, CuO and Cu(OAc)<sub>2</sub>  $\circ$  nH<sub>2</sub>O, preferably Cu, CuCl, Cu(OAc)<sub>2</sub>  $\circ$  nH<sub>2</sub>O, Cu(OH)<sub>2</sub>, Cu(OCH<sub>3</sub>)<sub>2</sub>, Cu<sub>2</sub>O and CuO, more preferably Cu, Cu(OH)<sub>2</sub>, Cu<sub>2</sub>O and CuO. In the above examples of the catalysts, n is normally an integer of 0 to 7. These catalysts may also be used as a mixture thereof, or may also be used after supported on heteropolyacids, silica gel, carbon powders, polymers, and other suitable carriers. The amount of the catalyst to be used is not particularly limited but is usually in the range of 0.01 – 120% by mole, preferably 0.1 – 10% by mole, relative to the olefin.

Examples of the aldehyde include formaldehyde, acetaldehyde, propionaldehyde, butanal, pentanal, hexanal, heptanal, decanal, 2 – methylpropanal, 2 – methylbutanal, cyclohexanecarboxaldehyde, isovaleral – dehyde, benzaldehyde, m – chlorobenzaldehyde, p – chlorobenzaldehyde, p – tolualdehyde, p – anisal – dehyde, and pivalaldehyde. When the oxidation is performed in the absence of a catalyst, 2 – methyl – propanal, 2 – methylbutanal, isovaleraldehyde, and pivalaldehyde are preferably used. The amount of the aldehyde to be used is not particularly limited but is usually in the range of 1 – 30 moles, preferably 1 – 10 moles, per mole of the olefin.

The addition of a proton source is particularly effective when the transition metal catalyst used is a simple substance. Examples of the proton source include formic acid, acetic acid, monochloroacetic acid, dichloroacetic acid, trichloroacetic acid, monobromoacetic acid, trifluoroacetic acid, propanoic acid, butyric acid, heptanoic acid, decanoic acid, benzoic acid, p-toluenesulfonic acid, hydrochloric acid, hydrogen bromide, sulfuric acid, nitric acid and water, preferably acetic acid and benzoic acid. The amount of the proton source to be used is not particularly limited, but is usually in the range of 1-100 moles per mole of the iron-containing or copper-containing catalyst. When the iron-containing or copper-containing catalyst contains water, however, the use of the proton source is not necessary.

In the process of the present invention, the reaction may also be performed in a solvent. Examples of the solvent which may be used include halogenated hydrocarbons such as dichloromethane, chloroform and ethylene dichloride, esters such as ethyl acetate, nitriles such as acetonitrile and aromatic hydrocarbons such as benzene, toluene, xylene, monochlorobenzene and dichlorobenzene.

The oxygen used in the present invention may be, besides oxygen, also air. The method of supplying the oxygen is not particularly restricted, and the reaction is generally conducted by blowing oxygen into the system, or under oxygen atmosphere, or supplying oxygen by other suitable means.

The method of feeding the olefin is not particularly limited but, when Fe<sub>2</sub>O<sub>3</sub> is used, the olefin is preferably fed last.

The reaction temperature is usually in the range from 0°C to the reflux temperature of the reaction mixture, preferably in the range from 20°C to 80°C.

The reaction time is not particularly limited. The reaction mixture may be analyzed by means of GC (gas chromatography) or the like, and the time at which the conversion to the intended epoxide levels off may be taken as the end point of the reaction. The reaction times is usually in the range from 1 to 48 hours.

In the present reaction, the aldehyde used is converted into the corresponding carboxylic acid and can be easily separated from the intended product.

After completion of the reaction, the intended epoxide can be obtained, for example, by recovering the catalyst by filtration and then subjecting the filtrate to washing with aqueous sodium hydrogencarbonate solution, then concentration and, if necessary, further operations such as rectification.

The process of the present invention is an industrially excellent one which can produce a corresponding epoxide from an olefin through a reaction with oxygen in the presence of an aldehyde, in the presence or absence of a proton source, and in the absence of a catalyst or by using an iron-containing or copper-containing catalyst which is easily available and recoverable and which brings little of undesirable substances into the waste water.

The present invention will be described in more detail below with reference to Examples, but it is in no way limited thereto.

### Example 1

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A mixture of 164 mg of cyclohexene, 433 mg of 2 – methylpropanal and 10 ml of dichloromethane was stirred overnight under an oxygen atmosphere at 25°C. Analysis of the reaction mixture by GC showed that cyclohexene oxide had been formed in 90% yield.

### Example 2

A mixture of 272 mg of  $\alpha$  – pinene, 517 mg of isovaleraldehyde and 10 ml of dichloromethane was stirred overnight under an oxygen atmosphere at 25 °C. Analysis of the reaction mixture by GC showed that  $\alpha$  – pinene oxide had been formed in 86% yield.

# Example 3

A mixture of 164 mg of cyclohexene, 517 mg of 2-metylbutanal and 10 ml of dichloromethane was stirred overnight under an oxygen atmosphere at 25°C. Analysis of the reaction mixture by GC showed that cyclohexene oxide had been formed in 88% yield.

# Examples 4 - 6

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Mixtures of 2 mmoles of an olefin, 517 mg of pivalaldehyde and 10 ml of dichloromethane were stirred overnight under an oxygen atmosphere at 25°C. Analysis of the reaction mixtures by GC gave the results shown in Table 1.

Table 1

Yield\*1) (%) Olefin Product Example No. 4 1 - Decene 1 - Decene oxide 65 5 trans - 5 - Decene trans - 5 - Decene oxide 92 6 cis - Stilbene oxide cis - Stilbene 38 trans - Stilbene oxide 48

Note:

\*1) Based on olefin

Examples 7 - 12

A solution of 914 mg of heptanal in 2 ml of dichloromethane was added dropwise over a period of 1 hour to respective mixtures of 164 mg of cyclohexene, 1% by mole (relative to cyclohexene) of a catalyst, 1.2 mg of acetic acid and 10 ml of dichloromethane under an oxygen atmosphere at 25°C, and the resulting mixtures were stirred for further 15 hours at the same temperature. Analysis of the reaction mixtures by GC gave the results shown in Table 2.

Table 2

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Example No. Cyclohexene oxide Catalyst yield\*1) (%) 7 Fe 71 8 FeCl<sub>2</sub> • 4H<sub>2</sub>O 6 FeSO<sub>4</sub> • 7H<sub>2</sub>O 9 18 10 FeCl<sub>3</sub> • 6H<sub>2</sub>O 11 Fe(OAc)<sub>3</sub> 30 12 10 Fe<sub>2</sub>O<sub>3</sub>

\*1) Based on cyclohexene

Examples 13 - 18

Mixtures of 281 mg of 1 - decene, 1% by mole (relative to 1 - decene) of a catalyst, 1.2 mg of acetic acid, 685 mg of heptanal and 10 ml of dichloromethane were stirred under an oxygen atmosphere at 25°C for 17 hours. Analysis of the reaction mixtures gave the results shown in Table 3.

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Table 3

Example No.	Catalyst	Conversion*1)	1 - Decene oxide yield*2) (%)
13	Fe	56	52(92)
14	FeCl <sub>2</sub> • 4H <sub>2</sub> O	11	5(49)
15	FeSO₄ • 7H <sub>2</sub> O	5	4(81)
16	FeO	6	5(84)
17	FeCl <sub>3</sub> • 6H <sub>2</sub> O	6	4(65)
18	Fe(OAc) <sub>3</sub>	9	7(82)

### Note:

- \*1) Based on 1 decene.
- \*2) Based on 1 decene. Values in parenthesis are based on converted 1 decene.

# Examples 19 - 24

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Mixtures of 281 mg of 1 – decene, 1.1 mg of Fe, 1.2 mg of acetic acid, 6 mmoles of an aldehyde and 10 ml of dichloromethane were stirred under an oxygen atmosphere at 25 °C for 17 hours. The reaction mixtures were analyzed by GC to obtain the results shown in Table 4.

# Table 4

Example No.	Aldehyde	Conversion*1) (%)	1 - Decene oxide yield*2) (%)
19	Acetaldehyde	3	3(100)
20	Butanal	73	59( 81)
21	2 - Methylpropanal	37	28( 75)
22	Cyclohexanecarboxaldehyde	30	25( 84)
23	Pivalaldehyde	44	32( 73)
24	Benzaldehyde	16	12( 77)

### Note:

- \*1) Based on 1 decene.
- \*2) Based on 1 decene. Values in parenthesis are based on converted 1 decene.

### Examples 25 - 29

Mixtures of 281 mg of 1 – decene, 1.1 mg of Fe, 1% by mole (relative to 1 – decene) of an acid, 685 mg of heptanal and 10 ml of dichloromethane were stirred under an oxygen atmosphere at 25°C for 17 hours. The reaction mixtures were analyzed by GC to obtain the results shown in Table 5.

Table 5

Example No. Acid Conversion\*1) (%) 1 - Decene oxide yield\*2) (%) 25 Benzoic acid 59 56(95) 26 Formic acid 56 51(91) 27 Heptanoic acid 58 53(91) 45(78) 28 Trifluoroacetic acid 58 29 p - toluenesulfonic acid 41 39(94)

10

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Note:

\*1) Based on 1 - decene.

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\*2) Based on 1 - decene. Values in parenthesis are based on converted 1 - decene.

# Examples 30 - 32

Mixtures of 281 mg of 1 – decene, 1.1 mg of Fe, 2.4 mg of benzoic acid, 685 mg of heptanal and 10 ml of a solvent were stirred under an oxygen atmosphere at 25°C for 17 hours. The reaction mixtures were analyzed by GC to obtain the results shown in Table 6.

Table 6

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Example No.	Solvent	conversion*1) (%)	1 - Decene oxide yield*2) (%)
30	Acetonitrile	61	56(92)
31	Benzene	31	31(99)
32	Ethyl acetate	12	12(99)

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Note:

\*1) Based on 1 - decene.

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\*2) Based on 1 - decene. Values in parenthesis are based on converted 1 - decene.

# Examples 33 - 34

Mixtures of 281 mg of 1-decene, 1.1 mg of Fe, 2.4 mg of benzoic acid, heptanal and 10 ml of dichloromethane were stirred under an oxygen atmosphere at 25°C for 17 hours. The reaction mixtures were analyzed by GC to obtain the results shown in Table 7.

Table 7

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Example No.	Heptanal equivalent (relative to 1 - decene)	Conversion*1) (%)	1 - Decene oxide yield*2) (%)
33	2	38	37(98)
34	4	64	55(86)

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Note:

\*1) Based on 1 - decene.

2) Based on 1 - decene. Values in parenthesis are based on converted 1 - decene.

# Examples 35 - 37

Mixtures of 281 mg of 1-decene, Fe, 2.4 mg of benzoic acid, 685 mg of heptanal and 10 ml of dichloromethane were stirred under an oxygen atmosphere at 25°C for 17 hours. The reaction mixtures were analyzed by GC to obtain the results shown in Table 8.

Table 8

Example No.	Fe equivalent (relative to 1-decene)	Conversion*1) (%)	1-Decene oxide yield*2) (%)
35	0.001	58	52(89)
36	0.05	56	48(85)
37	0.10	56	47(83)

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Note: \*1) Based on 1-decene.

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\*2) Based on 1-decene. Values in parenthesis are based on converted 1-decene.

### 30 Examples 38 - 45

Mixtures of 2 mmoles of an olefin, 1.1 mg of Fe, 2.4 mg of benzoic acid, 685 mg of heptanal and 10 ml of dichloromethane were stirred under an oxygen atmosphere at 25°C for 17 hours. The reaction mixtures were analyzed by GC to obtain the results shown in Table 9.

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Table 9

	Example No.	Olefin	Conversion*1) (%)	Product	Yield*2) (%)
40	38	Cyclohexene	100	Cyclohexene oxide	79( 79)
	39	trans - 5 - Decene	100	trans - 5 - Decene oxide	89( 89)
	40	cis - 5 - Decene	100	cis – 5 – Decene oxide trans – 5 – Decene oxide	56( 56) 30( 30)
45	41	1 - Methylcyclohexene	29	1 - Methylcyclohexene oxide	29(100)
	42	α - Pinene	23	α - Pinene oxide	23( 99)
	43	trans - Stilbene	14	trans - Stilbene oxide	12( 84)
50	44	cis - Stilbene	28	cis – Stilbene oxide trans – Stilbene oxide	5( 16) 21( 76)
	45	2 - Cyclohexenol	100	2,3 - Epoxy - 1 - cyclohexanol	38( 38)

Note:

\*1) Based on olefin

\*2) Based on olefin. Values in parenthesis are based on converted olefin.

# Examples 46 - 50

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Examples 38 - 45 were repeated except that 517 mg of pivalaldehyde was used in place of 685 mg of heptanal, to obtain the results shown in Table 10.

Table 10

Example No.	Olefin	Conversion*1) (%)	Product	Yield*2) (%)
46	1 - Methylcyclohexene	100	1 - Methylcyclohexene oxide	88(88)
47	α - Pinene	100	α - Pinene oxide	81(81)
48	trans - Stilbene	100	trans - Stilbene oxide	90(90)
49	cis – Stilbene	66	cis - Stilbene oxide trans - Stilbene oxide	9(13) 44(66)
50	Methyl p - methoxycinnamate	47	Methyl 3 - (4 - methoxyphenyl)glycidate	43(90)

#### Note:

- \*1) Based on olefin.
- \*2) Based on olefin. Values in parenthesis are based on converted olefin.

# 25 Examples 51 - 54

Mixtures of 272 mg of  $\alpha$  – pinene, 1.1 mg of Fe, 2.4 mg of benzoic acid, 6 mmoles of an aldehyde and 10 ml of dichloromethane were stirred under an oxygen atmosphere at 25 °C for 17 hours. The reaction mixtures were analyzed by GC to obtain the results shown in Table 11.

Table 11

Example No.	Aldehyde	Conversion*1) (%)	α - Pinene oxide Yield*2) (%)
51	Butanal	11	11(98)
52	Isovaleraldehyde	42	33(79)
53	2 - Methylpropanal	91	72(79)
54	Benzaldehyde	12	11(98)

# Note:

- \*1) Based on  $\alpha$  pinene.
- \*2) Based on  $\alpha$  pinene. Values in parenthesis are based on converted  $\alpha$  pinene.

# Example 55

A solution of 1.83 g of heptanal in 4 ml of dichloromethane was added dropwise over a period of 1 hour to a mixture of 561 mg of trans-5-decene, 2.2 mg of Fe, 2.4 mg of acetic acid and 20 ml of dichloromethane under an oxygen atmosphere at 25°C, and the resulting mixture was stirred for further 15 hours at the same temperature. The reaction mixture was analyzed by GC to obtain the following results.

Conversion: 87%

trans - 5 - Decene oxide yield :

84% (based on trans - 5 - decene)

97% (based on converted trans - 5 - decene)

cis - 5 - Decene oxide yield:

- 3% (based on trans 5 decene)
- 3% (based on converted trans 5 decene)

# Examples 56 - 66

Example

No.

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A mixture of 637 mg of benzaldehyde, 3.2 mg of  $Fe_2O_3$  and 12 ml of benzene was stirred vigorously under an oxygen atmosphere at room temperature, and then 2 mmoles of an olefin was added thereto over a period of 0.5 hours. After completion of the addition, the mixture was stirred for further 17 hours at the same temperature. The reaction mixture was analyzed by GC to obtain the results shown in Table 12.

Table 12

Yield\*2)

(Z)

97( 97)

90(90)

85(85)

81(81)

89(89)

96(100)

90(99)

100(100)

80(80)

84( 86)

88\*3

Product

Cyclohexene

α-Pinene oxide

Styrene oxide

styrene oxide

styrene oxide

1-Bromo-2.3-

Dichloro-2.3-

epoxybutane

Norbornene

α-Terpineol

oxide

oxide

1-(3,4-

Dimethoxy-

pheny1)-1,2-

5,6-Epoxy-3-

cholestanyl acetate

epoxy-propane

epoxybutane

trans-1,4-

p-Chloro-

p-Methoxy-

oxide

Conversion\*1)

100

100

100

100

100

96

91

100

100

98

(z)

Olefin

Cyclohexene

α-Pinene

Styrene

styrene

Crotyl

butene

bromide

trans-1.4-

Norbornene

 $\alpha$ -Terpineol

isoeugenol

Cholesteryl acetate

Methyl-

Dichloro-2-

p-Chloro-

p-Methoxystyrene

1	0

15

20

25

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35

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Note: \*1) Based on olefin.

- \*2) Based on olefin. Values is parenthesis are based on converted olefin.
- \*3) Isolated yield

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### Examples 67 - 71

A solution of 897 mg of cyclohexanecarboxaldehyde in 2 ml of dichloromethane was added dropwise over a period of 2 hours to a mixture of 164 mg of cyclohexene, 3% by mole (relative to cyclohexene) of a catalyst, 1.2 mg of acetic acid and 10 ml of dichloromethane under an oxygen atmosphere at 25°C, and the resulting mixture was stirred at the same temperature for further 15 hours. The reaction mixture was analyzed by GC to obtain the results shown in Table 13.

Table 13

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Example No. Catalyst Cyclohexene oxide yield\*1) (%) 67 Cu 80 68 Cu(OAc)<sub>2</sub> 85 69 CuCl 78 70 82  $Cu_2O$ 71 CuO 75

#### Note:

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### Examples 72 - 75

Mixtures of 281 mg of 1-decene, 1% by mole (relative to 1-decene) of a catalyst, 637 mg of benzaldehyde and 12 ml of dichloromethane were stirred under an oxygen atmosphere at 25°C for 17 hours. The reaction mixtures were analyzed by GC to obtain the results shown in Table 14.

Table 14

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Example No.	Catalyst	Conversion*1) (%)	1 - Decene oxide yield*2) (%)
72	Cu(OCH₃)₂	53	30(57)
73	Cu(OH)₂	51	32(64)
74	Cu	42	26(60)
75	Cu(OAc) <sub>2</sub>	40	25(63)

# Note:

- \*1) Based on 1 decene.
- \*2) Based on 1 decene. Values in parenthesis are based on converted 1 decene.

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# Examples 76 - 82

Mixtures of 2 mmoles of an olefin, 1.0 mg of Cu(OH)<sub>2</sub>, 673 mg of cyclohexanecarboxaldehyde and 12 ml of dichloromethane were stirred under an oxygen atmosphere for 17 hours. The reaction mixtures were analyzed by GC to obtain the results shown in Table 15.

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<sup>\*1)</sup> Based on cyclohexene.

Table 15

Example No.	Olefin	Conversion*1) (%)	Product	Yield*2) (%)
76	1 - Decene	51	1 - Decene oxide	27(83)
77	Cyclohexene	100	Cyclohexene oxide	79(79)
78	trans - 5 - Decene	83	trans - 5 - Decene oxide	80(96)
79	cis - 5 - Decene	96	cis – 5 – Decene oxide trans – 5 – Decene oxide	19(20) 63(66)
80	α – Pinene	100	α – Pinene oxide	84(84)
81	trans - Stilbene	93	trans - Stilbene oxide	76(82)
82	cis – Stilbene	81	cis - Stilbene oxide trans - Stilbene oxide	7( 9) 57(70)

Note:

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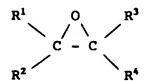
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\*1) Based on olefin.

\*2) Based on olefin. Values in parenthesis are based on converted olefin.

#### **Claims**

5 1. A process for producing an epoxide represented by the formula (2)



wherein  $R^1$ ,  $R^2$ ,  $R^3$  and  $R^4$  may be the same or different and each denotes a hydrogen atom,  $(C_1 - C_{20})$ alkyl group, alkyl group substituted with a halogen, hydroxy, alkoxy, phenoxy, acyloxy, acyl, alkoxycarbonyl or phenoxycarbonyl; phenyl group, phenyl group substituted with a halogen, alkyl, alkoxy, phenoxy, acyloxy, acyl, alkoxycarbonyl or phenoxycarbonyl; phenylalkyl group, phenylalkyl group substituted with a halogen, alkyl, alkoxy, phenoxy, acyloxy, acyl, alkoxycarbonyl or phenoxycarbonyl; acyl group, alkoxycarbonyl group or phenoxycarbonyl group; provided that  $R^1$  and  $R^2$  or  $R^1$  and  $R^3$  may combine with each other to form a ring, or  $R^1$ ,  $R^2$  and  $R^3$  may combine altogether to form a condensed ring, which comprises reacting an olefin represented by the formula (1)

$$C = C$$

$$R^{3}$$

$$R^{2}$$

wherein R1, R2, R3 and R4 are the same as defined above, with oxygen in the presence of an aldehyde.

2. A process for producing an epoxide represented by the formula (2) which comprises reacting an olefin represented by the formula (1) with oxygen in the presence of an aldehyde, in the presence or absence of a proton source and in the presence of an iron – containing catalyst or a copper – containing catalyst.

- 3. The process according to Claim 1, wherein the aldehyde is 2-methylpropanal, 2-methylbutanal, isovaleraldehyde or pivalaldehyde.
- The process according to Claim 2, wherein the iron containing catalyst is Fe, FeCl₂ nH₂O, FeSO₄ nH₂O, FeCl₃ nH₂O, Fe(OAc)₃ or Fe₂O₃.
  - 5. The process according to Claim 2, wherein the iron containing catalyst is Fe or Fe<sub>2</sub>O<sub>3</sub>.
- 6. The process according to Claim 2, wherein the copper-containing catalyst is Cu, CuCl, Cu(OAc) 2 nH<sub>2</sub>O, Cu(OH)<sub>2</sub>, Cu(OCH<sub>3</sub>)<sub>2</sub>, Cu<sub>2</sub>O or CuO.
  - 7. The process according to Claim 2, wherein the copper-containing catalyst is Cu, Cu(OH)2, Cu2O or CuO.
- 15 8. The process according to Claim 2, wherein the proton source is acetic acid or benzoic acid.
  - 9. The process according to Claim 2, wherein the amount of proton source is 1 100 moles per mole of the iron containing or copper containing catalyst.
- 20 10. The process according to Claim 1 or 2, wherein the amount of the aldehyde is 1 30 moles per mole of the olefin.
  - 11. The process according to Claim 1 or 2, wherein the amount of the catalyst is 0.01 120% by mole relative to the olefin.
  - 12. The process according to Claim 1 or 2, wherein the reaction is performed in a solvent.

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- 13. The process according to Claim 12, wherein the solvent is dichloromethane, ethyl acetate, acetonitrile or benzene.
- 14. The process according to Claim 1 or 2, wherein the reaction temperature is 0°C to the reflux temperature of the reaction mixture.
- 15. The process according to Claim 1 or 2, wherein the reaction time is 1 to 48 hours.
- 16. The process according to Claim 1 or 2, wherein the oxygen is supplied by blowing it into the reaction system.

# **EUROPEAN SEARCH REPORT**

Application Number

EP 92 11 8554 Page 1

ategory	Citation of document with it of relevant pa	ndication, where appropriate,	Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int. CL5)
	•	F AKTIENGESELLSCHAFT) t, particularly page 2, lines 11, 12 and	1,3,10, 12-16	C07D301/06 C07D303/02
<b>x</b>	examples 1-3 * FR-A-1 367 771 (IMPERIAL CHEMICAL INDUSTRIES LIMITED) * the whole document, particularly page 2, left-hand column, paragraph 4 and examples 2 and 3 *			
	FR-A-1 323 787 (IMPERIAL CHEMICAL INDUSTRIES LIMITED)  * the whole document, particularly example 11 and page 7, table VI, runs 3 and 7 *		1-16	
	FR-A-1 377 981 (SHAWINIGAN CHEMICALS IMITED) the whole document, particularly examples 10 and 11 *		1-16	TECHNICAL FIELDS SEARCHED (bt. Cl.5)
	R-A-1 400 305 (EASTMAN KODAK COMPANY) the whole document *		1,3,10, 12-16	C07D
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	DE-A-1 937 387 (BASF AKTIENGESELLSCHAFT)  * the whole document, particularly the examples *		1,3,10, 12-16	
(	DE-A-1 568 407 (DEU WISSENSCHAFTEN ZU B * the whole documen	ERLIN)	1,3,10, 12-16	
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	The present search report has h			
T	THE HAGUE Data of completion of the search  O4 FEBRUARY 1993			ALLARD M.S.

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A: technological background
O: non-written disclosure
P: intermediate document

å : member of the same patent family, corresponding document



# **EUROPEAN SEARCH REPORT**

Application Number

EP 92 11 8554 Page 2

	lication, where appropriate,	to claim	CLASSIFICATION OF THE APPLICATION (Int. CL5)	
	ITUT FRANÇAIS DU NTS ET LUBRIFIANTS)	1-16		
LIMITED)		1-16		
			TECHNICAL FIELDS SEARCHED (Int. Cl.5)	
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The present search report has be				
Place of search THE HAGUE	Date of completion of the search O4 FEBRUARY 1993		ALLARD M.S.	
erticularly relevant if taken alone erticularly relevant if combined with and scument of the same category	E : earlier patent di after the filing o ther D : document cited L : document cited	T: theory or principle underlying the invention E: earlier patent document, but published on, or after the filling date D: document cited in the application L: document cited for other reasons		
	* the whole document BE-A-682 335 (IMPERI LIMITED)  * thw whole document examples 1 and 3 *  The present search report has be Place of search THE HAGUE  CATEGORY OF CITED DOCUMENT ricularly relevant if taken alone ricularly relevant if combined with ano	The present search report has been drawn up for all claims  The present search report has been drawn up for all claims  Place of search THE HAGUE  CATEGORY OF CITED DOCUMENTS  Pricularly relevant if taken alone riticularly relevant if combined with another cument of the same category theological background  The whole document deed to the search part of	* the whole document *  BE-A-682 335 (IMPERIAL CHEMICAL INDUSTRIES LIMITED)  * thw whole document, particularly examples 1 and 3 *   The present search report has been drawn up for all claims  Pisce of search THE HAGUE  CATEGORY OF CITED DOCUMENTS Tricularly relevant if cambined with another cument of the same category theological background  Ti theory or principle underlying the strictling relevant if cambined with another cument of the same category Theological background  T: theory or principle underlying the strictling relevant if cambined with another cument of the same category Theological background  T: theory or principle underlying the strictling gate  D: document deed in the application of the same at	